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EXHIBIT 1

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Biological Response Modifiers

Joel.W. Goldwein, MD, Brad Somer, MD, and the Oncokink Team Abramson Cancer Center of the University of Pennsylvania Last Modified: November 1, 2001

Introduction

Biological response modifiers (BRMs) are another form of chemotherapy sometimes administered to cancer patients. They modify the relationship between the tumor and the patient by strengthening the patient's blological response to tumor cells. BRMs can be divided into three major categories eccording to mechanism of action:

- agents that restore, augment, or modulate the patient's normal immunological mechanisms;
- agents that have direct antitumor effects; and
- 31. agents that have other blologic effects, such as interference with a fumor cell's ability to metastasize or survive after metastasis, promotic of cell differentiation, or interference with neoplastic transformation in cells.

Scientists began studying BRMs in cancer therapy in the 1960s, labeling the treatment modality immunotherapy. After promising results in animal studies researchers militated many large-scale clinical trials to stimulate cancer. patients. Immune systems using the bacterial agents dacillus Calmette-Gueri (BCG) and Corynebacterium parvum (C. parvum). The results of these trials were discouraging, so the research into immunotherapy as a possible modali for cancer treatment lost momentum.

Medical

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Recent advances have prompted a renewed interest in BRMs, and today biological response modification is an important area in cancer research and treatment.

OncoLink **Art Gallery** Confronting Caricer

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Through Art is an exhibition by people whose lives have been muched by cancer.



Today's featured works: by Bruce Pollock

Immune System: Background

The body's limitude system mounts a coordinated combination of nonspecific The body's immune system mounts a coordinated comordation in independent and specific responses to foreign substances (e.g. microbes; and certain othe toxins, called antigens). Both physical injury and the presence of antigens ca invoke nonspecific host defenses: These defenses include physical barriers ar chemical factors, such as the skip and mucous membranes, addic gastric, secretions, and normal intestinal flora. The "inflammatory response" is anothinospecific host defense that serves to control the growth of microorganisms. and prevent systemic infection.

Specific immune responses are ellofted by the presence of an antigen. These reactions are characterized by a memory: following the initial exposure to an antigen, specific portions of the limmune system produce memory cells that entigen, specific portains of the infiltratic system product a more victorius response to subsequent exposures to the same antigen. These specific memory responses are generally divided into humora: and cell-mediated immunity.

Humoral Immunity refers to the immunity conferred by the B-lymphocyte cel produced by the lymph system. These lymphocytes, also colled the B-cells, produce antibodies. Antibodies are small proteins that can deactivate antigen by a variety of mechanisms, usually by binding with them. Antibody antigen interaction is specific. Only one type of antibody can interact and meutralize at the contract and meutralize at t interaction is specific. Unity one type or antipody, can interact and mentifices specific type of antipea. This interaction then activates the "complement cascade," a system of proteins that "complements" antibody activity by destroying bacteria and helping the body rid itself of antibody/antigen complexes.

Cell-mediated immunity raters to the immunity conferred by the mutation of Cell-mediated immunity refers to the immunity conferred by the mutation of lymphocytes, which is thought to occur in the thyrius gland. These lymphocytes, also called *T cells*, directly or indirectly destroy viruses, malignant cells, cells infected with intracellular organisms, and cells of grafte organs. Different types of Ticells have different immune functions: cytotoxic cells directly destroy and gens, helper Ticells activate the "humoral immune system" and cytotoxic Ticells; and suppressor Ticells inhibit antibody production and other immune responses.

Other cells that are important in the immune response are macrophages and natural killer (NK) cells. Macrophages are white blood cells with a number of Important functions. They bind to an antigen and "present" the antigen to undifferentiated cells (precursor cells); these, in turn, become activated and undirerendatau cella threcolor cells / triose, in thin, become available produce mature lymphocytes. Without this macrophage processing, the T and B cells could not respond to some types of antigens. NK cells are cytotoxic to tumor cells and virus-infected cells.

Many cells in the immune system produce chemicals that aid in regulating the immune response. These substances are referred to as mediators and broad! referred to as cytokines. Many cytokines are under study, to determine their effection the immune systems.

Types of BRM Therapy

A brief review of BRM agents currently being evaluated follows.

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Romocional Authorates

The use of monocional antibodies (MoAbs) involves the development of speciantibodies directed against antigens located on the surface of tumor cells.

Samples of the patients tumor calls are taken and processed to reveal specif antipodes to the turbor associated antigens. In order for this approach to work, a sufficient quantity of antigens unique to the turbor associated antigens unique to the turbor cells must be present. In addition, the turbor antigens must be sufficiently different from the turbor antigens must be sufficiently different from the turbor of the turbor antigens must be sufficiently different from the turbor of turbor of the turbor of turbor of the turbor of turbor of the turbor of tur present. In addition, the turnor another to provoke an antibody response.

The antibodies can be used either alone to Idli cancer cells or as carriers of other substances used for either therefreute or diagnostic purposes. For example, chemotherapeutic agents can be attached to monadonal antibodies to deliver high concentrations of these toxic substances directly to the tumor cells. In theory, this approach is less toxic and more effective that conventional interpretations are required to the conventional chemotherapy because the returns the delivery of harmful agents. conventional chemotherapy because it reduces the delivery of harmful agents to normal tissues is decreased:

Monoclonal antibodies can also be used for diagnostic purposes. They may be used to carry radioactive substances to cancer cells, thus pinpointing the location of metastases previously undetected by other methods.

Despite these uses, some monoclonal antibodies have limitations. Because some monoclonal antibodies may be made using mouse antibodies, they are, themselves; foreign proteins that often trigger an immune response; thus, they can be neutralized before any therapeutic effect occurs. In addition, monocionals may lack specificity for tumor antigens. Tumor cell antigens may not be different enough from those on normal cells to ensure only cancer cell destruction; studies have revealed that most monoclonal antibodies interact with antigens on both normal and cancer cells.

More recently, many monoclonal antibodies have been created which are only derived from human proteins. Some are already FDA-approved and many derived from human proteins. Some are already FDA-approved and many others are in clinical trials, with approval imminent. In general, they have proven useful in treatment of memotologic malligrancies and lymphoma. In addition, monoclopals are in development for use against solid rumors. All of these antibodies have multiple potential applications including nuclear imagin surgical mapping, and direct therapy in multiple settings (alone, in conjunctic with chemotherapy, for treatment of memotises, in adjuvant settings, in high dose rates; etc.) In the future this field will most likely grow in importance in the war against cancer.

In the clinical setting, the apeutic monoclonal antibodies are usually given av-4-6 hours by continuous intravenous Infusion. Because of the risk of serious: allergic:reactions (particularly with the mouse antibodies), patients are premedicated with acetaminophen and an antihistaminie and monitored closely. Emergency drugs are kept by the bedside: Potential side effects of monocional antibodies include dyspical and mild wheezing fever, chills, headache, rasis, mausea, vomitino; tactiveantia, and alleggic reactions. headache, rash, mausea, vomiting, tachycardia, and allergic reactions.

Research studies are currently underway using monoclonals for a variety of diseases; include T cell lymphoma, chronic and acute lymphocytic leukemia. melanoma, colorectal cancer, and neuroblastoma.

Interferons .

Interferons (IFNs) are small proteins that inhibit viral replication and promote

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problement is desirable to OncoLink | The Web's First Cancer Resource

the cellular (Ticell) immune response. Interferon use for cancer treatment w. himited until the late 1970s, when technological advances enabled mass production of IFN.

There are currently three major types of IFNst alpha, beta, and gamma. Each type has similar but distinctive capabilities for altering biological responses.

Alpha-IFN was the first BRM approved by the Food and Drug Administration (FDA) in 1986, two different manufacturers have braints of this product available: Its main indication is for use in treatment of heavy to the its currently also indicated for use in the treatment of harrycell leukemia and currently also indicated for use in the treatment of harrycell leukemia and ADS-associated Kaposi's surcoma. It has also defricins taked therapeutic effectiveness against hematologic diseases such as low-grade Hodgkin's effective on some myelogenous leukemia. It has also proven to be some what effective on some myelogenous leukemia against hematologic diseases against hematologic diseases such as low-grade Hodgkin's effective on some myelogenous leukemia against hematologic diseases against hematologic diseases such as low-grade Hodgkin's effective on some leukemia against hematologic diseases and against hematologic diseases against hematologic diseases

Interferons may produce side effects of varying frequency and intensity depending on dose, schedule, route of administration, and the type of IFN. There is currently a "ance per week" formulation of INF in late clinical trials which reduces the overall side-effects expendenced by patients. One of the which reduces the overall side-effects expendenced by patients. One of the which reduces the overall side-effects expendenced by patients. One of the which reduces the correlation side effects of IFN therapy is a flu-like syndrome. Symptoms include fever, chills, tachycardia, muscle aches, malaise, fatigue, and headaches. This reaction is extremely common during a patient's first exposure to IFN, but usually decreases in intensity with continued therapy.

Other common side effects to IFN include a decreased white blood cell count, anemia (with prolonged the apply), and decreased platelets: Gastrointestinal symptoms such as a loss of appetite, nauses, womiting, and diarrhea may als be present. Central nervous system toxicities range from mild confusion and sleepiness to selzures. Acute vidney failure is rare, but can occur, loss of hal may also be a problem.

Interferon can be administered by IV bolus of Infusion, or Intramuscular, subcutaneous, or intrattrecal injection. It can also be given thranasally. Redness and initiation at the injection site may occur. Since IFN is often administered on an outpatient basis, it is essential that the patient and family are rought the technique of administration and how to manage side effects. are taught the technique of administration and how to manage side effects.

Interleukin-2
Interleukin-2 (II-2) is a substance produced by lymphocytes. In addition to Interleukin-2 (II-2) is a substance produced by lymphocytes. In addition to being an essential factor for the growth of Ticells, II-2 addition to Fellow II-2 also activates lymphocytes. In addition II-2 also activates lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated with II-2. Lak cells destroy tumor cells and lymphocytes are incubated in additional incubated responses to II-2 therapy.

The most severe toxicities result from IL-2's ability to increase capillary permeability. This may cause hypotension, ascites, generalized body edema, and pulmonary edema.

Chilis and fever also frequently occur within a few hours after IL-2.

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administration. Headache, malaise, and other fullike symptoms are also common. Gastrointestinal effects include nausea, vomiting, loss of appetite, darrhea, and mucositis. Some liver dysfunction is common during therapy by resolves once treatment; is stopped. Central nervous system toxicity if nesolves once treatment; is stopped. Central nervous system toxicity if nesolves once treatment; is stopped. Central nervous system toxicity if nesolves once treatment; is stopped. Central nervous system toxicity if nesolves in confusion, and consider in an interesting in the kidneys is and consultation, and consultation, and course in the consultation, and mala, and a decrease in plateless are more likely with higher cumulative, doses. Skin changes include redness, rash, pruritus, and occasionally skin desquamation.

Atthough many research studies with II-2 require intensive supportive care is acute care settings, other comment regimens can be given on an acute care settings, other contract the strength of the settings of the contract outpatient basis. Patient addication in these structures is especially important outpatient basis. because patients must be alert to potential side effects that should be report.

Colony Stimulating Factors

Colony Stimulating Factors

Colony Stimulating factors (CSFs) are growth factors whith mediate the proliferation, maturation; regulation, and activation of granulocytes, many macrophage, lymphocytes, monocytes; erythrocytes, and platelets. Many macrophage, lymphocytes, monocytes; erythrocytes, and platelets. Many macrophage, lymphocytes, monocytes; erythrocytes, and platelets. Many use and some are in various stages of clinical trials. Generally, CSFs have be named for the major cell lineage they affect. Granulocyte macrophage CSF (GM-CSF) affects both granulocytes and macrophage illneage; granulocyte CSI (GM-CSF) targets both granulocytes. These two have been FDA-approved. The main indication is for treatment of neutropenic fevers. This has ensured multiple scenarios, including the prevention of neutropenic fevers primarily o secondarily, and for use in stem cell mobilization. Other colony stimulating factors include pleutopletin IL-3, or multiple CSF, which affects early cell factors include pleutopletin IL-3, or multiple CSF, which affects early cell lineages; and macrophage CSF (M-CSF) targets macrophage production. Neumega is an IL-11 that indices platelet growth (and has FDA approval) an was hoped to limit the amounts of platelet transfusions patients may require as originally hoped, and therefore is not often used. Other colony stimulating factors include thrombopoetin and platelet-derived growth factor (POGF), factors include thrombopoetin and platelet-derived growth factor (POGF), factors include the many facturers to strongly consider removing from the prompting their manufacturers to strongly consider removing from the market. Exphroporatin, which targets exphirocyte production, was approved the PDA in 1989 for use in anemia caused by end-stage-renal disease (Epo (Im)). Another version, manufactured by Ortho Blotech (Procrit) is used to treat anemia; related to cancer and cancer thereby as well and the fatigue

GM-CSF and G-CSF have been administered by TV bolus; subcutaneously by daily injection, or by continuous TV infusion. G-CSF therapy has been associated with only minimal toxicity, mainly bone pain. GM-CSF produces associated with only minimal toxicity, mainly bone pain. GM-CSF produces associated with only minimal toxicity. The daily follows the continuous and ambigurantees are according to the continuous and ambigurantees are -inore systemic containes, manually rangue, revers masse action, assertes ma and diarrhea. Blood levels of alkaline phosphatase and aminotransferases ma also be increased.

Medical use of these growth factors is an important step in understanding an manipulating the immune system. Their efficacy in the treatment of congenit hematologic diseases and their ability to reduce neutropedia during cancer

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treatment, makes them important agents in the treatment armamentarium.

Tumor Necrosis Factor Tumor necrosis factor (TNF) is a substance naturally secreted by macrophages. When activated by sindotoxias, the macrophages release TNF, which then binds to receptors on cell membranes. Once bound to the cell membrane, TNF initiates cellular activity and is possibly cytotoxic to that cell.

TNF is in the early phases of clinical trials and has not yet demonstrated therapeutic effectiveness against mallgnant diseases. Side effects of TNF are similar to those experienced with interferon therapy, including a flu-like syndrome and soreness at the injection site. Fevers and chills are generally mild and disappear with subsequent doses of TNF.

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